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## (54) Title: POLYNITRILE OXIDES

## (57) Abstract

The present invention is an approas dispersion of a stable polysistific cubic represented by the structure: R-(CaN-O<sub>3s</sub>, when x is an interpretate than 1, it is an arranks, sliptack, or opticallylatic group having a test one substanced subjects to each strike oxide cubic cubic proposed and statement support. The present invention is too a model and object mechanisms of the statement content in the cubic propose and statemented groups. The present invention is not a model and story water from the nature. The present invention is not provided a simple matter of presenting expense creating passes that the cubic cubic cubic proposed as strained in the nature. The present invention is more constituting the steps of a planting the latter with a stable polysistic cubic, and it summaring value from the nature. The present invention to make the cubic cubic cubic cubic provides a simple matter of presenting expenses.

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# POLYNITRILE OXIDES

The present invention relates to stable polynitrile oxides.

Nitrile oxides react with unsaturated compounds to form cyclic compounds. For example, nitrile oxides react with a) olefins and alkynes to form loxazolines and loxazoles, respectively; b) aldehydes and ketones to form 1,3.4-dioxazoles; c) this canonyls to form thioxazoles; d) imino compounds to form 1,2.4-oxadiazolines; e) isocyanates to fo rm 1,2.4-oxadiazolinones; and f) carboxyls to form hydroximic acids. (See Grundmann and Grunanger, Mirit Oxides, Springer, New York, pp. 5-319, 1971.)

Nitrile oxides an be prepared by a number of methods, most notably from the dehydrohalogenation of the corresponding hydroximic add halide, which can be prepared by the halogenation of the corresponding aldoxime. The aldoxime, in turn, can be prepared by reacting the corresponding aldoxyde with a hydroxyl amine. General methods that teach the preparation of ritille loxides are described in hylitic plotes, surar, po. 3 Fides. surar, po. 3 Fides.

15 Because Intrile oxides tend to dimerzie in the absence of stabilizing groups, it is desirable to either prepare the intille oxides in gain, or to prepare stabilized mixtue invalue. Nitrile oxides can be stabilized by the presence of substituents, such as ethyl, methyl, or methylsulfide groups adjacent to the nitrile oxide group (see Nitrile Oxides, puzze, p. 14). Examples of their initial initial exides, including stable bis-intrile oxides are disclosed in Nitrile.

20 Oxides, supra, pp. 16-21; [xv. Akad. Nauk SSSB, Ser. Khim., No. 5, pp. 1201-1203 (1991); and [xv. Akad. Nauk SSSB, Ser. Khim., No. 7, pp. 1609-1615 (1991)] One such stable bis-nitrile oxide, 2.4.6-triethylberaene-1,3-bis(nitrile oxide), has been shown to be useful for the vulcanization of natural ninber

None of the above-cited art suggests the use of stable polynitrile oxides as 25 curatives for latexes. It would be an advance in the art to cure latexes using a one-part, room-

The present invention is a water-insoluble aqueous dispersion comprising a stable polynitrile oxide represented by the structure:

G-(C=N-O')<sub>x</sub>

temperature curative

30 where x is an integer greater than 1, G is an aromatic, aliphatic, or cycloaliphatic group having at least one substituent adjacent to each intile oxide group, the substituent characterized by inhibiting dimirization of nitrile oxide, and being non-interfering with a reaction between nitrile oxide groups and unsaturated groups.

In a second aspect, the present invention is a method of curing a latex having a 35 polyunsaturated disperse phase, comprising the steps of:

> a) mixing with the latex a water-insoluble, stable polynitrile oxide represented by the structure:

> > G-(C=N-O:).

where x is an integer greater than 1, preferably an integer from 2 to 6, G is an aromatic, alighattic, or cycloalighatic group having at least one substituent adjacent to each nitrile oxide group, the substituent characterized by inhibiting dimerization of nitrile oxide, and being non-interfering with a reaction between nitrile oxide groups and unsaturated groups; and b) removing water from the mixture, preferably by evaporation.

In a further aspect, the present invention is a compound having the structure:



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15 wherein each if is independently C<sub>T</sub>CT2=likyl, F, Cl, Br., (I, O,C,C<sub>T</sub>-2-likyl, or S,C<sub>T</sub>-C<sub>12</sub>-alkyl; each R<sup>4</sup> is a substituent that does not spontaneously react with the nitrile exide group; each if is independently 0, 1, or 2; n<sup>4</sup> is an integer greater than 1; each X! is independently a bond or a connecting group; and Y is a polyvalent radical containing an ether, ester, amide, amine, carbonate, lettone, urethane, anylene, or thioether molety; or each X' and Y' together are a pond connecting the bereater enlique the betareare in the property of the proper

The present invention provides a simple means of preparing one-part coating systems that can be cured at room temperature without the release of by-products.

The polynitrile oxides suitable for the practice of the present invention are

hindered polynitrile oxides. The term "polynitrile oxide" is used herein to refer to two or more aromatic nitrile oxide groups per molecule. It is to be understood that the term "aromatic" includes heteroaromatic moleties such as pyridines, funare and thiophenes. The term "unsaturated" is used herein to denote a late of the type A = A,' or A = A,' where A is a carbon atom, and A' is a carbon, oxygen, nitrogen, suffur, or phosphorus atom. For the purpose of this invention, a nitrile oxide group is not an unsaturated group. The term "polyunsaturated" is used herein to denote more than one unsaturated group. The preferred unsaturated group include oleffice and alignes.

Each nitrile oxide is adjacent to at least one substituent that is 1) unreactive with nitrile oxide and 2) non-interfering with the reaction between the nitrile oxide groups and unsaturated groups, preferably olefinically or acetylenically unsaturated groups.

Traditionally, nitrile oxides are prepared in situ in the presence of an unsaturated substrate with which the nitrile oxides are intended to react. However, the stable polynitrile oxide used as a curing agent in the present invention can be prepared separately and is sufficiently stable in the absence of the reactive substrate to be effective as a curing agent.

Preferably, the stable polynitrile oxide forms less than 10 percent, more preferably less than 5 percent, and most preferably less than 1 percent dimers in 30 days at room temperature. Examples of hindered aromatic polynitrile oxides include:

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$$(^{-}O-N^{+}=C)_{m}$$
 $(^{-}C=N^{+}-O^{-})_{n}$ 
 $(^{-}C=N^{+}-O^{-})_{n}$ 
 $(^{-}C=N^{+}-O^{-})_{n}$ 
 $(^{-}C=N^{+}-O^{-})_{n}$ 
 $(^{-}C=N^{+}-O^{-})_{n}$ 

$$\mathbb{R}^{1}$$
 $\mathbb{R}^{1}$ 
 $\mathbb{R}^{1}$ 
 $\mathbb{R}^{3}$ 
 $\mathbb{R}^{3}$ 

where R1, R2, and R3, and R4 are each independently H, R, halo, SH, SR, SD, SD, R, hydroxy, or OR, with the proviso that at least one of R1, R2, R3, and R4 that is adjacent to a nitrille isotice group is north; R3, R6, R7, and R4 are each independently H, R, halo, SH, SH, SR, SD, SD, R, Morroy, or OR, wherein R is a C₁-C1₂ linear, branched, or cyclic ably group, preferably a C₁-C4, 25 linear or branched allyl group, more preferably ethyl or methyl; or R4 and R3, or R7 and R4. together with the carbon atoms to which they are attached, form a benzare ring, wherein at least one of R7 or R7 in not H, and at least one of R6 or R8 in not H; ils 2 or 3; m and n are each 0, 1, or 2, and n + m ≥ 2, preferably 2 or 3.

Other examples of hindered aromatic polynitrile oxides include compounds represented by the following structures:

$$(^{\circ}0-N^{+}=C)_{L}$$
 $R^{9}$ 
 $R^{10}$ 
 $(C=N^{+}-0^{-})_{R}$ 
 $R^{12}$ 

where R<sup>9</sup>, R<sup>10</sup>, R<sup>11</sup>, and R<sup>12</sup> are each independently H, R, halo, SH, SR, SOR, SO<sub>2</sub>R, hydroxy, or OR with the proviso that at least one of R<sup>9</sup> and R<sup>11</sup> is not H when a nitrile oxide group is adjacent to

-3-

(~0-N\* = C) p R11

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both  $\mathbb{R}^3$  and  $\mathbb{R}^{11}$ , and at least one of  $\mathbb{R}^{10}$  and  $\mathbb{R}^{12}$  is not  $\mathbb{N}$  when a nitrille oxide group is adjacent to both  $\mathbb{R}^{10}$  and  $\mathbb{R}^{12}$ ;  $\mathbb{N}$ ,  $\mathbb{$ 

and

Examples of specific hindered aromatic polynitrile oxides that are suitable for the practice of the present invention include the following compounds:

$$C = N_{+} - 0$$
.

 $C = N_{+} - 0$ .

 $C = N_{+} - 0$ .

 $C = N_{+} - 0$ .

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Stable aliphatic or cycloaliphatic polynitrile oxides can be prepared from a suitably functionalized aliphatic or cycloaliphatic polyaldehyde. The polyaldehyde can then be reacted with hydroxylamine to form the polyaldoxime, which can then be treated with bleach and caustic treatment to form the desired aliphatic polynitrile oxide.

A suitably functionalized aromatic mononitrile oxide or monoaldehyde can be used to prepare a polynitrile oxide represented by the following formula:

wherein each R' is independently  $C_1$ — $C_2$ - $C_3$ - $C_3$ ,  $C_3$ ,  $C_4$ - $C_4$ - $C_4$ - $C_4$ - $C_3$ - $C_3$ - $C_4$ - $C_4$ - $C_4$ - $C_3$ - $C_4$ 

Suitably functionalized hindered aromatic mononitrile oxides or monoaldehydes preferably indude 2,6-disubstituted benzonitrile oxides or benzaldehydes having an ester, acetate, hydrox, epoxy, fluorine, chlorine, bromine, or doine group commeted directly to the benzene ring or indirectly through a connecting group. Preferably, the suitably functionalized 2,6-disubstituted benzonitrile oxide or benzaldehydes is represented by the following structure:

where R', R', X', and n' are previously defined; Q is -C≡N\*O- or -CHO; and Z' is an ester, actate, amine, hydroxy, epoxy, amide, keto, aldehyde, fluorine, chlorine, bromine, or iodine 1s group.

For example, 3-hydroxymethyl-2.4.6-trimethylenzonitrile oxide or its corresponding benzaldehyde precursor can be: (a) transseterified with a diester or condensed with a diadd chloride to form a dintrile oxide diester; (b) reacted with phospene to form a dintrile oxide containing a carbonate group; (d) reacted with a discoyanate to form a dintrile oxide oxide containing urethane groups; (d) reacted with a dibenzyl chloride to form a dintrile oxide containing two ether groups; (e) reacted with a diglycldyl ether to form a dintrile oxide containing two ether groups; (e) reacted with a diglycldyl ether to form a dintrile oxide oxide and the discovery of the discove

Similarly, the suitably functionalized hindered aromatic nitrile oxide can be 25 reacted with a second suitably functionalized hindered aromatic nitrile oxide to form a dinitrile oxide. For example, 3-hydroxymethy-2,45-trimethylbenzonitrile oxide on be reacted with 3-chloromethyl-2,6-dimethylbenzene nitrile oxide to form a bisfinitrile oxide) dimethyl ether.

Polynkrile oxides having a functionality of greater than 2 (for example, a trinitrile oxide) can readily be prepared by reacting a dinkrile oxide with a compound having more than 30 2 unsaturated siste. For example, 2,4,6-triethylbenzene-1,3-dinkrile oxide can be reacted with trimeturated propane trisorylate to form the following trinitrile oxide:

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An aqueous dispersion of the stable polymitrile oxide is prepared, then advantageously combined with an aqueous dispersion of a polymasturated monomer or polymer or a combination thereof, to make a stable multicomponent dispersion. The term "stable multicomponent dispersion" is used herein to mean that microscopic mixing (and 5 therefore, the reaction rate) of the polymitrile oxide and the polyuristurated monomer and/or polymer is slower than it would be in the absence of the aqueous medium. Preferably, the extent of the reaction between the polymitrile oxide and polyurasturated monomer and/or polymer dispersions is less than 10 percent in 8 hours, more preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most percent in 30 days, and most preferably less than 10 percent in 30 days, and most preferably less than 10 percent in 30 days, and most percent in 30 days.

10 The aqueous dispersion of the polynitrile oxide can be prepared by emulsifying an emulsifiable concentrate of the polynitrile oxide. This concentrate can be prepared, for example, by mixing a solution of the polynitrile oxide with a surfactant.

The polynitrile oxide may itself be prepared as a surfactant, for example, by reacting an excess of a dinitrile oxide with a polyunsaturated surfactant;

or, for example, by reacting a trinitrile oxide with a monounsaturated surfactant:

where X<sup>e</sup> is a hydrophilic group, such as a poly(oxyethylene), a carboxylate, or a sulfate.

Other methods of forming a polynitrile oxide surfactant include reacting the polynitrile oxide with a polymeric surfactant having polyunsaturation:

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